







## Statistica Computazionale Progredito

Campionamento Hamiltonian Monte Carlo (HMC) con il software Stan

Leonardo Egidi (legidi@units.it)

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What is Stan? • Why Stan? • Writing a Stan program • Linked package: bayesplot

## Origins



Stanislaw Ulam (1909-1984): Manhattan project, H-Bomb experiments in Los Alamos, MCMC father jointly with John von Neumann.

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- Writing a Stan program
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#### What is Stan?

- Probabilistic programming language and inference algorithms.
- Stan program
  - declares data and (constrained) parameter variables
  - defines log posterior (or penalized likelihood)
- Stan inference
  - MCMC for full Bayes
  - Variational Bayes for approximate Bayes
  - Optimization for (penalized) MLE
- Stan ecosystem
  - lang, math library (C++)
  - interfaces and tools (R, Python, Julia, many more)
  - documentation (example model repo, user guide & reference manual),
     case studies , R package vignettes)
  - online community ( Stan Forums on Discourse)

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# Why Stan?

- Fit rich Bayesian statistical models. Close to the big data philosophy.
- Efficiency
  - Hamiltonian Monte Carlo + NUTS
  - Compiled to C++
- Flexible domain specific language
- "Freedom-respecting, open-source"
  - doc & written materials
  - interacting community
  - continuous development
- Interaction with some other R packages designed to explore the Stan output.

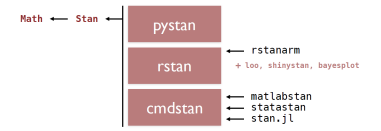
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## Who is using Stan?

- Biological & physical sciences: clinical trials, epidemiology, genomics, population ecology, entomology, ophthalmology, neurology, agriculture, fisheries, cancer biology, astrophysics & cosmology, molecular biology, oceanography, climatology.
- Social sciences: population dynamics. psycholinguistics, social networks, political science, human development, economics.
- Many more: sports analytics, public health, publishing, finance, pharma, actuarial, recommender systems, educational testing, materials engineering.

#### Interfaces

# Interfaces + Tools



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## Improving MCMC performance

With Stan, we aim to provide an MCMC implementation that works robustly for as many target distributions as possible

- Gibbs, RW Metropoilis can be very inefficient, hard to diagnose.
- To explore complicated high-dimensional spaces we need to leverage what we know about the geometry of the typical set.
- For such a reason, Stan enjoys Hamiltonian Monte Carlo.

The Stan users may use, analyze and interpret HMC outputs as they were standard MCMC outputs.

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## Before starting

#### What is a Bayesian model?

- Building a Bayesian model forces us to build a model for how the data is generated
- We often think of this as specifying a prior and a likelihood, as if these are two separate things
- They are not!

#### Generative models

The philosophy behind Stan is to think generatively.

The model is expressed as a joint probability distribution of observed and unobserved variables, which may be decomposed as follows:

$$p(y,\theta) = p(y|\theta)\pi(\theta) \tag{1}$$

The posterior of interest is then proportional to the joint distribution (1):

$$p(\theta|y) \propto p(y|\theta)\pi(\theta)$$
 (2)

### Generative models

A Bayesian modeller commits to to an a priori joint distribution:

$$p(y,\theta) = \underbrace{p(y|\theta)\pi(\theta)}_{\textit{Likelihood} \times \textit{Prior}} = \underbrace{\pi(\theta|y)p(y)}_{\textit{Posterior} \times \textit{Marginal Likelihood}} \tag{3}$$

## Generative models and vague priors

#### What is the problem with *vague/diffuse* priors?

- If we use an improper prior, then we do not specify a joint model for our data and parameters.
- More importantly, we do not specify a data generating mechanism p(y).
- By construction, these priors do not regularize inferences, which is quite often a bad idea
- Proper but diffuse is better than .improper but is still often problematic.

## Generative models

- If we disallow improper priors, then Bayesian modeling is generative.
- In particular, we have a simple way to simulate from p(y):



## Stan computations

Stan works in logarithmic terms: all the computations are actually done on log-scale. So, for the posterior we have.

$$\log(\pi(\theta|y)) = \log(\pi(\theta)) + \log(p(y|\theta)) + \text{constant}$$
 (4)

Products become sums of logs:

$$p(y|\theta) = \prod_{i=1}^n p(y_i|\theta) \rightarrow \log(p(y|\theta)) = \sum_{i=1}^n \log(p(y_i|\theta)).$$

## Starting point

We are now going to write a Stan program together:

- Open a new empty file in RStudio
- Save it as linear\_regression.stan

## Blocks strategy

#### Stan programs are organized into blocks:

- data block: declare data types, sizes, and constraints. Read from data source and constraints validated. Evaluated: once.
- parameters block: declare parameter types, sizes, and constraints. Evaluated: every log prob evaluation.
- transformed parameters block: declare those parameters transformed from the original ones declared in the parameters block. Evaluated: every log prob evaluation.
- model block: statements defining the posterior density in log scale. Evaluated: every log prob evaluation.
- generated quantities: declare and define derived variables. (P)RNGs, predictions, event probabilities, decision making. Constraints validated. Evaluated: once per draw.

## Data block

```
data {
 // Dimensions
 int<lower=1> N;
 int<lower=1> K;
 // Variables
 matrix[N, K] X;
 vector[N] y;
```

```
/* multiple lines of
comments */
```

## Parameters' block

```
parameters {
  real alpha;
  vector[K] beta;
  real<lower=0> sigma;
```

constraints required in parameters block

#### Model block

```
model {
    // priors (flat, uniform, if omitted)
    sigma ~ exponential(1);
    alpha ~ normal(0, 10);
    for (k in 1:K) beta[k] ~ normal(0, 5);

    for (n in 1:N) {
        y[n] ~ normal(X[n, ] * beta + alpha, sigma);
    }
}
```

Why is the default automatically uniform?

- $\pi(\theta) \propto 1$  (0 on log scale)
- Nothing added to log prob

## Generated quantities block

```
generated quantities {
  vector[N] y_rep;
  for (n in 1:N) {
    real y_hat = X[n,] * beta + alpha; // local/temp
    y_rep[n] = normal_rng(y_hat, sigma);
  }
}
```

## Complete Stan model

```
data {
  int<lower=1> N:
  int<lower=1> K:
  matrix[N, K] X;
  vector[N] v:
parameters {
  real alpha:
  vector[K] beta;
  real<lower=0> sigma;
model {
  sigma \sim exponential(1);
  alpha \sim normal(0, 10);
  for (k \text{ in } 1:K) \text{ beta}[k] \sim \text{normal}(0, 5):
  for (n in 1:N)
    y[n] ~ normal(alpha + X[n, ] * beta, sigma);
generated quantities {
  vector[N] y rep;
  for (n in 1:N)
    y_rep[n] = normal_rng(alpha + X[n,] * beta, sigma);
```

Observed variables

Unobserved variables

 $\log \pi(\theta)$  $\log p(y | \theta)$ 

Simulate from generative model

## Launching the Stan model from R

Now we may launch the Stan program directly in R:

```
library(rstan)
# passing the data (already stored)
data <- list(N=N, K=K, X=X, y=y)
# fitting the model
fit1 <- stan(
  file = 'linear_regression.stan',
  data = data.
  iter = 2000,
  chains = 4)
# extracting the estimates
sims <- extract(fit1)</pre>
```

## First example: 8 schools

This example studied coaching effects from eight schools.

We denote with  $y_{ii}$  the result of the *i*-th test in the *j*-th school. We assume the following model:

$$y_{ij} \sim \mathcal{N}(\theta_j, \sigma_y^2)$$
  
 $\theta_j \sim \mathcal{N}(\mu, \tau^2)$ 

Do some schools perform better/worse according to these coaching effects? Here is the data, already aggregated by schools:

schools\_dat <- list(J = 8,  

$$y = c(28, 8, -3, 7, -1, 1, 18, 12),$$
  
 $sigma = c(15, 10, 16, 11, 9, 11, 10, 18))$ 

```
// saved as 8schools.stan
data {
 int<lower=0> J; // number of schools
 real y[J];
             // estimated treatment effects
 real<lower=0> sigma[J]; // standard error of effect estimates
parameters {
 real mu;
                      // population treatment effect
 real<lower=0> tau; // standard deviation in treatment effects
 vector[J] eta;
               // unscaled deviation from mu by school
transformed parameters {
 vector[J] theta = mu + tau * eta;  // school treatment effects
model {
 eta \sim normal(0,1);
                    // prior
 y ~ normal(theta, sigma); //likelihood
```

## First example: 8 schools

To fit the model and visualize the estimates, it is sufficient to type in R the following commands (with 2000 iterations and 4 chains as a default):

```
fit_8schools <- stan(file = '8schools.stan', data = schools_dat)
print(fit_8schools, pars=c("mu", "tau", "theta"))</pre>
```

```
2.5% 25% 50% 75% 97.5% n_eff Rhat
               sd
         mean
                               7.92 11.05 18.05
         7.89 5.04 -2.31 4.74
                                                2352
                                                      1
mu
tau
         6.70 5.71 0.24 2.61
                               5.43 9.19 21.16 1480
                                                      1
theta[1] 11.36 8.23 -2.25 6.18 10.29 15.46 31.15
                                                3161
theta[2] 7.89 6.21 -4.43 3.96 7.83 11.78 20.47
                                                4923
                                                      1
theta[3]
                               6.56 10.81 20.25
        6.05 7.59 -10.81 1.92
                                                4057
                                                      1
theta[4] 7.60 6.44 -5.36 3.74
                               7.63 11.73 20.57
                                                5055
                                                      1
theta[5] 5.13 6.23 -8.45 1.35
                                                4346
                                                      1
                               5.60 9.26 16.37
theta[6] 5.95 6.68 -8.21 1.99
                               6.30 10.21 18.21
                                                4313
                                                      1
theta[7]
        10.62 6.93 -1.58 6.12 10.14 14.53 25.63
                                                3381
                                                      1
theta[8]
         8.40 7.77 -7.18 3.84 8.26 12.63 25.78
                                                3854
                                                      1
```

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# Posterior graphical analysis with bayesplot

Once we fit a model, it is to vital check it via graphical inspection. The bayesplot package (for any help, see the vignette) is designed to this task.

The package allows to display:

- Posterior uncertainty intervals
- Univariate marginal posterior distributions
- Bivariate plots
- Trace plots
- Posterior predictive plots

library(bayesplot)

## Posterior graphical analysis with bayesplot

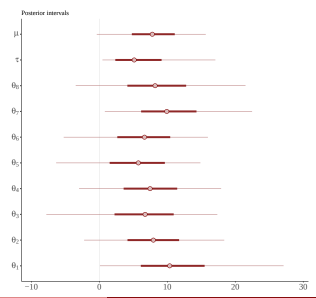
The first step is to save the posterior. Then you have many choices:

```
posterior <- as.array(fit_8schools)</pre>
mcmc_intervals(posterior)
                                # posterior intervals
mcmc_areas(posterior)
                                # posterior areas
```

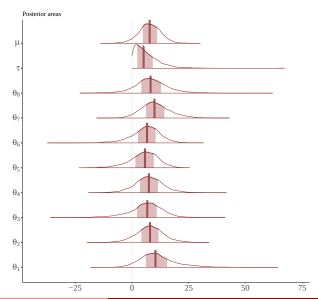
mcmc\_dens(posterior) # marginal posteriors mcmc\_pairs(posterior) # bivariate plots mcmc\_trace(posterior) # trace plots

With the arguments pars or regex\_pars you may select the desired parameters.

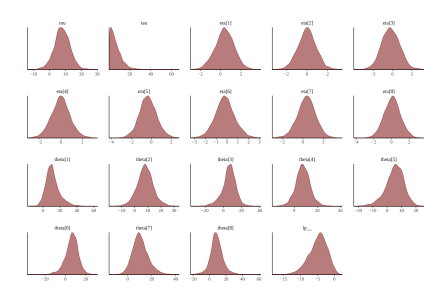
## Posterior uncertainty intervals



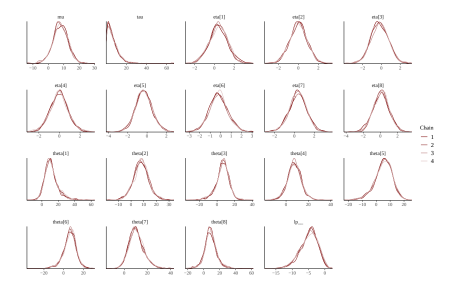
# Posterior uncertainty areas



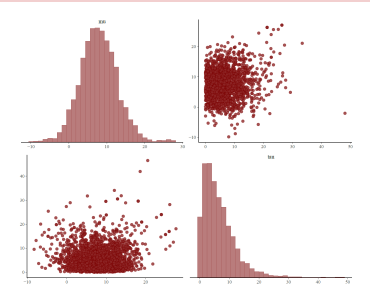
## Marginal posteriors



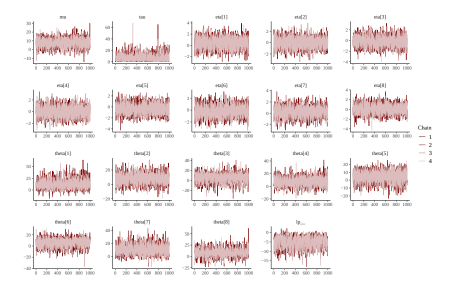
## Marginal posteriors separated for each chain



# Bivariate posterior plots



## Trace plots for the Markov chains



## Our challenge with Stan

The Stan shuttle is ready to start! We will learn to:

- write simple and more complex model in Stan: Im, glm, hierarchical models.
- analyze the posterior summaries.
- criticize the model and, eventually, change/reparametrize it.

# Further reading

#### Further reading:

- Gelman, A., et al. (2013). Bayesian Data Analysis, 3rd edition, CRC Press, Boca Raton (Section 12.6)
- Carpenter, B, and Gelman, A, Hoffman, M.D., Lee, D., Goodrich, B., Betancourt, M., Brubaker, M., Guo, J., Li, P., Riddell, A. (2017). Stan: A Probabilistic Programming Language, Journal of statistical software 76(1). Here the  $\bigcirc$